

ERDEY, Laszlo, prof., dr. (Budapest, XI., Gellert ter 4); SVEHLA,
Gyula, dr. (Budapest, XI., Gellert ter 4)

Accuracy of silver determination by atomic absorption
methods. Acta chimica Hung 41 no.1/2:187-194 '64.

1. Institute of General Chemistry of Budapest Technical
University.

ERDEY, László, prof. dr. (Budapest, XI., Gellert ter 4); LIPTAY, György, dr.
(Budapest, XI., Gellert ter 4); DAVID, Péter (Budapest, II., Lovcház u.39)

Derivatographic study of thermal decomposition of electrical insulating materials and insulators. Periodica polytechnica electr 8 no. 3:242-250 '64.

1. Department for General Chemistry of the Polytechnical University, Budapest, and Research Institute for Electrical Industry, Budapest.
2. Editorial Board Member, "Periodica Polytechnica - Electrical Engineering" (for Erdey). Submitted February 10, 1964.

ERDEY, Laszlo, prof., dr. (Budapest, XI., Gellert ter 4); KASA, Imre, dr. (Budapest, XI., Gellert ter 4)

Examination of 2-hydroxy-4-amino-4'-methoxy-diphenylamine redox indicator. Acta chimica Hung 41 no.1/2:59-65 '64.

1. Institut fur Allgemeine Chemie der Technischen Universitat Budapest. 2. Mitglied, Redaktionskollegium, "Acta Chimica Academiae Scientiarum Hungaricae" (for Erdey).

LIPTAY, Gyorgy, dr okleveles vegyeszmernok, adjunktus; DAVID, Peter, okleveles vegyesz; ERDEY, Laszlo, dr., okleveles vegyesz, akademikus

Derivatographic analysis of the heat caused decomposition of electric insulators and insulations. Pt.1. Elektrotechnika 57 no.9:392-397 S '64.

1. Chair of General Chemistry, Budapest Technical University, Budapest XI., Gellert ter 3 (for Liptay). 2. Research Institute of Electric Industry, Budapest, VI., Népkoztársaság útja 32 (for David). 3. Head, Chair of General Chemistry, Budapest Technical University, Budapest XI., Gellert ter 3 (for Erdey).

L 63740-65 ENT(1)/IJP(c)

ACCESSION NR: AT5021739

HU/2502/64/041/01-/0037/0042

AUTHOR: Erdely, László (Erdel, L.) (Doctor, Professor) (Budapest); Buzas, Ilona (Buzash, I.) (Doctor) (Budapest); Takacs, József (Takach, Y.) (Budapest)

TITLE: Contribution to the luminescence mechanism of lucigenine

SOURCE: Academia scientiarum hungaricae. Acta chimica, v. 41, no. 1-2, 1964, 37-42

TOPIC TAGS: luminescence, light emission, laboratory optic instrument, catalysis

ABSTRACT: [German article] The luminescence of lucigenine was investigated under various experimental conditions with the aid of a modified Magnephot I microlumen meter. The lighting mechanism was found to be composed of two components, one reversible and the other irreversible. Catalysts such as ethyl alcohol, butyl alcohol, and osmium tetroxide, caused a shift in the ratio of these two components towards the irreversible. The emission of light without the presence of catalysts coincided with the formation of minute quantities of N-methylacridone.

Orig. art. has: 3 graphs.

Card 1/2

L 63740-65

ACCESSION NR: AT5021739

ASSOCIATION: Institut für Allgemeine Chemie der Technischen Universität, Budapest
(Institute for General Chemistry at the Technical University)

SUBMITTED: 03May63

ENCL: 00

SUB CODE: OP, GO

NR REF SOV: 001

OTHER: 004

JPRS

Card

2/2

L 63681-65

ACCESSION NR: AT5021741

HU/2502/64/041/01-/0059/0065

AUTHOR: Erdey, Laszlo (Erdel, L.) (Professor, Doctor) (Budapest); Kasa, Imre
(Kasha, I.) (Doctor) (Budapest) S
B+1

TITLE: Investigation of the oxidation-reduction indicator 2-hydroxy-4-amino-
4 prime-methoxy-diphenylamine

SOURCE: Academia scientiarum hungaricae. Acta chimica, v. 41, no. 1-2, 1964,
59-65

TOPIC TAGS: oxidation reduction reaction, diphenylamine

ABSTRACT: [German article] The transition potential of the indicator, the number of electrons participating in the oxidation-reduction mechanism, the reversibility of the indicator mechanism, and the absorption spectrum of the indicator were established and discussed. Two protons and two electrons participate in the oxidation process which yields violet-colored p-quinoidal compounds; no semiquinone was detected. The

Card 1/2

L 63681-65

ACCESSION NR: AT5021741

indicator performs reversibly in the 2-12 pH range; it is suitable within these limits for mildly oxidizing systems. Orig. art. has: 2 tables, 5 graphs, 2 formulas.

ASSOCIATION: Institut für Allgemeine Chemie der Technischen Universität, Budapest
(Institute for General Chemistry at the Technical University)

SUBMITTED: 22May63

ENCL: 00

SUB CODE: CC, CC

NR REF SOV: 000

OTHER: 009

JPRS

llc
Card 2/2

63677-65

ACCESSION NR: AT5021747

HU/2502/64/041/01-/0109/0122

AUTHOR: Erdey, Laszlo (Erdei, L.)(Doctor, Professor)(Budapest); Paulik, Ferenc
Buzach-Gere, Eva (Buzag, E.)Budapest); Polos, Laszlo (Polosh, L.)

TITLE: Derivatographic and electron-microscopic examination of barium sulfate precipitates. Part 2

SOURCE: Academia scientiarum hungaricae. Acta chimica, v. 41, no. 1-2, 1964, 109-122

TOPIC TAGS: chemical precipitation, barium compound, sulfate, electron microscopy

ABSTRACT: Barium sulfate precipitates obtained in various analytical precipitations were examined by derivatography and electron microscopy. Pure barium sulfate was obtained only from very dilute solutions even after all volatile impurities were eliminated by calcination. Eighteen electron micrographs and 9 derivatographic curves were presented and discussed. Orig. art. has: 27 figures, 1 table.

ASSOCIATION: Institut fur allgemeine Chemie der Technischen Universitat, Budapest (Institute for General Chemistry, Technical University)

Card 1/2

L 63677-65

ACCESSION NR: AT5021747

SUBMITTED: 03Jan64

ENCL: 00

SUB CODE: GC, OP

NR REF SOV: 001

OTHER: 020

JPRS

llc
Card 2/2

L 63899-65 EPF(c)/EWP(j) RM

ACCESSION NR: AT5022529

HU/2502/64/042/002/0131/0144

AUTHOR: Csuros, Zoltan (Chyuryesh, Z.) (Professor, Doctor) (Budapest); Dusa, Zsigmond (Dusa, Zh.) (Budapest); Petro, Jozsef (Petro, Y.) (Doctor) (Budapest); Erdely, Laszlo (Erdely, L.) (Professor, Doctor) (Budapest); Paulik, Ferenc (Budapest)

TITLE: Investigations on catalysts. Part 40: Investigations on Raney-nickel catalysts. Section 15: Effects of the alkali used as extractant and of the hydrogen content on the activity

SOURCE: Academiae scientiarum hungaricae. Acta chimica, v. 42, no. 2, 1964, 131-144

TOPIC TAGS: nickel, catalysis, hydrogen, basic catalysis

ABSTRACT: A derivatographic method was developed for the study of pyrophoric catalysts such as those from Raney-nickel. The method was applied to catalysts prepared by using various solvents such as sodium hydroxide, potassium hydroxide, and sodium carbonate solutions. Catalysts prepared by using KOH or NaOH contained relatively high quantities of hydrogen and the hydrogen content was in proportion to their nickel content. However, no relation was evident between the catalyst's composition and its effectiveness. Orig. art. has 1 graph and 4 tables.

Card 1/2

L 63899-65

ACCESSION NR: AT5022529

ASSOCIATION: Institute of Organic Chemical Technology, Technical University,
Budapest; Institute of General Chemistry, Technical University, Budapest

SUBMITTED: 10Feb64

ENCL: 00

SUB CODE: GC

NO REF SOV: 002

OTHER: 012

JPRS

llc
Card 2/2

KOCSIS, Elemér; RÖT, László

Examination of the adaptability of measuring devices to spectrometers. Gep 12 no.1 2-4 Jan 1965.

1. Chair of General Chemistry of Budapest Technical University.

INCZÉDY, János, dr. (Budapest, XI., Gellert ter 4); NEMESHEGYI, Gabor (Budapest, XI., Gellert ter 4); ERDEY, László, prof., dr. (Budapest, XI., Gellert ter 4)

Separation and determination of rare earth metals by ion exchange chromatography. Pts.1-2. Acta chimica Hung 43 no.1:1-15 '65.

1. Institute of General Chemistry of Budapest Technical University.
Submitted July 2, 1964.

L 63187-65

ACCESSION NR: AT5021755

HU/2502/64/041/01-/0187/0194

AUTHOR: Khalifa, H. (Kalifa, Kh.)(Doctor)(Giza); Erdex, László (Erdei, L.)(Doctor, Professor)(Budapest); Svehla, Gyula (Shvekhla, D.)(Doctor)(Budapest)

TITLE: Accuracy of silver determinations by atomic absorption methods 7
B+

SOURCE: Academia scientiarum hungaricae. Acta chimica, v. 41, no.1-2, 1964, 187-194

TOPIC TAGS: silver, spectrophotometry, chemical detection

ABSTRACT: The optimum experimental conditions for the determination of silver by atomic-absorption spectrophotometry were determined and various calibration curves were presented. Foreign metals did not interfere with the determinations. The errors of the determinations in various concentration ranges (varying from 10 to 1300 p.p.m.) were established and presented in tables. Orig.art. has: 7 tables, 2 figures, 1 formula.

ASSOCIATION: Institute of General Chemistry, Technical University, Budapest

SUBMITTED: 22Jan64

NR REF SOV: 000

ENCL: 00

OTHER: 004

SUB CODE: IC,NP

JPRS

mll
Card 1/1

L 63899-65 EPF(c)/ENP(j) RM

ACCESSION NR: AT5022529

HU/2502/64/042/002/0131/0144

AUTHOR: Csuros, Zoltan (Chyuryash, Z.) (Professor, Doctor) (Budapest); Dusza, Zsigmond (Dusa, Zh.) (Budapest); Petro, Jozsef (Petro, Y.) (Doctor) (Budapest); Erdey, Laszlo (Erdei, L.) (Professor, Doctor) (Budapest); Paulik, Ferenc (Budapest)

TITLE: Investigations on catalysts. Part 40: Investigations on Raney-nickel catalysts / Section 15: Effects of the alkali used as extractant and of the hydrogen content on the activity

SOURCE: Academiae scientiarum hungaricae. Acta chimica, v. 42, no. 2, 1964, 131-144

TOPIC TAGS: nickel, catalysis, hydrogen, basic catalysis

ABSTRACT: A derivatographic method was developed for the study of pyrophoric catalysts such as those from Raney-nickel. The method was applied to catalysts prepared by using various solvents such as sodium hydroxide, potassium hydroxide, and sodium carbonate solutions. Catalysts prepared by using KOH or NaOH contained relatively high quantities of hydrogen and the hydrogen content was in proportion to their nickel content. However, no relation was evident between the catalyst's composition and its effectiveness. Orig. art. has 1 graph and 4 tables.

Card 1/2

L 63899-55

ACCESSION NR: AT5022529

ASSOCIATION: Institute of Organic Chemical Technology, Technical University,
Budapest; Institute of General Chemistry, Technical University, Budapest

SUBMITTED: 10Feb64

ENCL: 00

SUB CODE: GC

NO REF SOV: 002

OTHER: 012

JPRS

llc
Card 2/2

L 1179-66

ACCESSION NR: AT5025201

HU/2502/64/042/004/0379/0382

AUTHOR: Liptay, Gyorgy (Doctor)(Budapest); Hegyaljai Kias, Geza (Doctor)(Budapest);
Erdey, Laszlo (Professor, Doctor)(Budapest).

TITLE: Investigation by thermal analysis of the pyrolytic dehydrogenation of sterols

SOURCE: Academia scientiarum hungaricae. Acta chimica, v. 42, no. 4, 1964, 379-382

TOPIC TAGS: thermal analysis, pyrolysis, dehydrogenation, alcohol

Abstract: [English article] The pyrolytic decomposition of $\Delta^1,4$ -androsta-
diene-3,17-dione and of $\Delta^1,4,6$ -androstratriene-3,17-dione was investiga-
ted by thermal analysis employing the Orion GYEM 676 type derivatograph.
The curves obtained indicated that the splitting temperature of the angular
methyl group is not affected by the presence of the unsaturated B-ring, and
the first-mentioned compound pyrolyzed at a higher exothermic rate.
Orig. art. has 4 formulas and 2 figures.

ASSOCIATION: Department of General Chemistry, Technical University, Budapest;
Chinoin Factory of Pharmaceutical and Chemical Products, Budapest

SUBMITTED: 12 May 64

NO. REF SOV: 000

Card 1/1

ENCL: 00

OTHER: 008

SUB CODE: 00, 00

JPRS

L 41682-66 EWP(t)/ETI IJP(c) JD/JG

ACC NR: AT6031101

SOURCE CODE: HU/2502/65/043/002/0095/0100

AUTHOR: Erdey, Laszlo--Erdei, L. (Professor; Doctor); Kasa, Imre--Kasha, I. (Doctor); Kovacs, Laszlo--Kovach, L.

ORG: Technical University of Budapest, Institute of General Chemistry; Frederic Joliot-Curie Institute of Radiation Biology, Budapest

TITLE: Investigation of the thermoluminescent properties of lithium fluoride

SOURCE: Academia scientiarum Hungaricae. Acta chimica, v. 43, no. 2, 1965, 95-100

TOPIC TAGS: ^{v1}lithium fluoride, thermoluminescence

ABSTRACT: Changes in the thermoluminescent properties of lithium fluoride which take place on the effect of different factors were investigated. Thermoluminescence was found to depend on the nature of the lithium compound which served as an initial substance for the preparation of lithium fluoride. The same physical influences had different effects on lithium fluoride preparations obtained from various starting materials. Thermoluminescence was markedly increased by the addition of calcium fluoride. The authors thank Grad.-Engr. O. Roka for construction of the measuring device and for assistance with the measurements. Thanks are also given to Grad.-Engr. E. Kocsis for the spectrographic analysis. Orig. art. has: 4 figures and 1 table. /JPRS: 33,540/

SUB CODE: 07, 20 / SUBM DATE: 20Feb65 / ORIG REF:001 / SOV REF:001 / OTH REF:014

Card 1/1 af

0918 2321

L 47233-66 LIP(c)
ACC NR: AF6034307

SOURCE CODE: HU/0005/66/000/006/0268/0269

AUTHOR: Erdey, Laszlo; Kantor, Tibor 37B

ORG: Academic Research Group of Technical Analysis, Department of General Chemistry,
Technical University, Budapest (Muszaki Egyetem, Altalanos-Kemial Tanszek, Muszaki
Analitikai Akademiai Kutato Csoprt)

TITLE: Continuous introduction of powdered substances into spectroscopic light sources

SOURCE: Magyar kemial folyoirat, no. 6, 1966, 268-269

TOPIC TAGS: spectroscopy, spectroscopic analysis

ABSTRACT: A device is described which can be used for the continuous introduction of solid, powdered materials into arc and spark light sources. The substance is introduced through a tube-electrode with the aid of a crew spindle which is rotated at a constant speed by an electric motor. The "tube-electrode method" is simple and versatile, and can be applied in various spectroscopic analyses. Orig. art. has: 1 figure. [JPRS: 36,862]

SUB CODE: 20 / SUBM DATE: 26Sep65 / ORIG REF: 002 / OTH REF: 010

Card 1/1 hs

ERDEYNE SCHNEER, Anna

Some newer methods for rock and mineral analysis. Magy kem lap 19 no.6:
325-329 Je '64.

1. Research Group of Inorganic Chemistry, Hungarian Academy of
Sciences, Budapest.

ERDEYNE SCHNEER, Anna, a kémiai tudományok kandidátusa

Newest results in inorganic chemical qualitative analysis.
Kém tud közl MTA 22 no.1:71-88 '64.

1. Chair of General Chemistry, Lorand Eotvos University,
Budapest.

RADNOT, Magda; VEYNSHTEYN, P.[Weinstein, P.], doktor med.nauk, nauchnyy red.;
CHAPODI I.[Csapodi, I], doktor med. nauk, nauchnyy red.; SIZA,
Mario[Sziza, Mario, translator]; ERDI, K., otv. red.; CHERGE, I.
[Csorgo, I.], tekhn. red.

[Atlas of eye diseases]Atlas glaznykh boleznei. Budapest, Akade-
miai Kiado. Vol.2. 1963. 199 p. (MIRA 15:12)

1. Chlen-korrespondent Akademii nauk Vengrii.
(EYE—DISEASES AND DEFECTS)

ERDEY-GRUZ, Tibor; MAJTHENYI, Lajos

Migration mechanism of hydrogen and hydroxyl ions. Pt. 5.
Magy kem folyoir 65 no. 5:167-174 My '59.

1. Eotvos Lorand Tudomanyegyetem Fizikai-Kemiai Tanszeke,
Budapest.
2. "Magyar Kemiai Folyoirat" felelos szerkesztoje (for Erdey-
Gruz).

ERDEY-GRUZ, Tibor; MAJTHENYI, Lajos

Migration mechanism of hydrogen and hydroxyl ions. Pt. 6.
Magy kem folyoir 65 no. 6:212-218 Je '59.

1. Eotvos Lorand Tudomanyegyetem Fizikai-Kemai Tanszek, Budapest.
2. "Magyar Kemai Folyoirat" felelos szerkesztoje (for Erdey-Gruz).

ERDEY-GRUZ, Tibor, dr., akadémikus, egyetemi tanár (Budapest)

The fight and unity of the opposites in thermodynamics. Term tud
kozl 5 no.2:65-66 F '61.

ERDEY-GRUZ, Tibor

In commemoration of the 250th anniversary of the birth of Mikhail
Vasil'evich Lomonosov. Magyar kem lap 16 no.12:529-530 D '61.

ERDEY-GRUZ, T. (Budapest); DEVAY, J. (Budapest)

Raising the depolarisation of quicksilver electrodes
by alternating current. Rev chimie 7 no. 1: 181-188
'62.

1. Lehrstuhl für physikalische Chemie und Radiologie der
Roland-Eotvos-Universität; Elektrochemische Forschungs-
gruppe der Ungarischen Akademie der Wissenschaften,
Budapest.

ERDEY-GRUZ, Tibor, dr.

Answers by Dr. Tibor Erdey-Grus, president of the Council of Science and Higher Education. *Műszaki és Tudományok* 17 no.24:3 22 N '62.

1. Tudományok és Felsőoktatási Tanács elnöke.

ERDEY-GRUZ, Tibor; DEVAY, Jozsef; VAJASDY, Irma

Effect on sine currents on electrode processes.X. Effect of
sine currents on the hydrogen overvoltage of platinum cathode.
Magy kem folyoir 68 no.5:185-190 My '62.

1. Eotvos Lorand Tudomanyegyetem Fizikai-Kemiai es Radiologiai
Tanszeke, Budapest, es Magyar Tudomanyos Akademia Elektrokemiai
Kutato Csoportja, Budapest. 2. "Magyar Kemiai Folyoirat"
felelos szerkesztoje (for Erdey-Gruz).

ERDEY-GRUZ, Tibor; DEVAY, Josef; SZEGEDI, Robert

Effect of sine currents on electrode processes. XI. Effect of alternating currents on the Hg-Zn corrosion in the case of the processes of mixed control. Magyar kem folyoir 68 no.5:190-193 My '62.

1. Eotvos Lorand Tudomanyegyetem Fizikai-Kemiai Tanszeka, Budapest, es a Magyar Tudomanyos Akademia Elektrokemiai Kutato Csoportja, Budapest. 2. "Magyar Kemiai Folyoirat" felelos szerkesztoje (for Erdey-Gruz).

ERDEY-GRUZ, Tibor; DEVAY, Jozsef; HORANYI, Gyorgy; VAJASDY, Irma

The effect of sinusoidal current on electrode processes.XII.
Magy kem folyoir 68 no.9:373-376 S '62.

1. Eotvos Lorand Tudomanyegyetem Fizikai-Kemiai es Radiologiai
Tanszeke, Budapest, es Magyar Tudomanyos Akademia Elektrokemiai
Kutato Csoportja. 2. "Magyar Kemiai Folyoirat" felelos
szerkesztoje (for Erdey-Gruz).

SZABO, Zoltan, egyetemi tanar; POLINSZKY, Karoly, a kémiai tudományok doktora; MATOLCSY, Kalman, a kémiai tudományok kandidátusa; LEVAY, Gyula; NAGY, Ferenc, a kémiai tudományok doktora; BERECS, Endre, a kémiai tudományok kandidátusa docens; KORACH, Mor, ~~akadémikus~~; LENGYEL, Sandor, a kémiai tudományok doktora; SCHAY, Geza, akadémikus, egyetemi tanar; ERDEY-GRUZ, Tibor, akadémikus

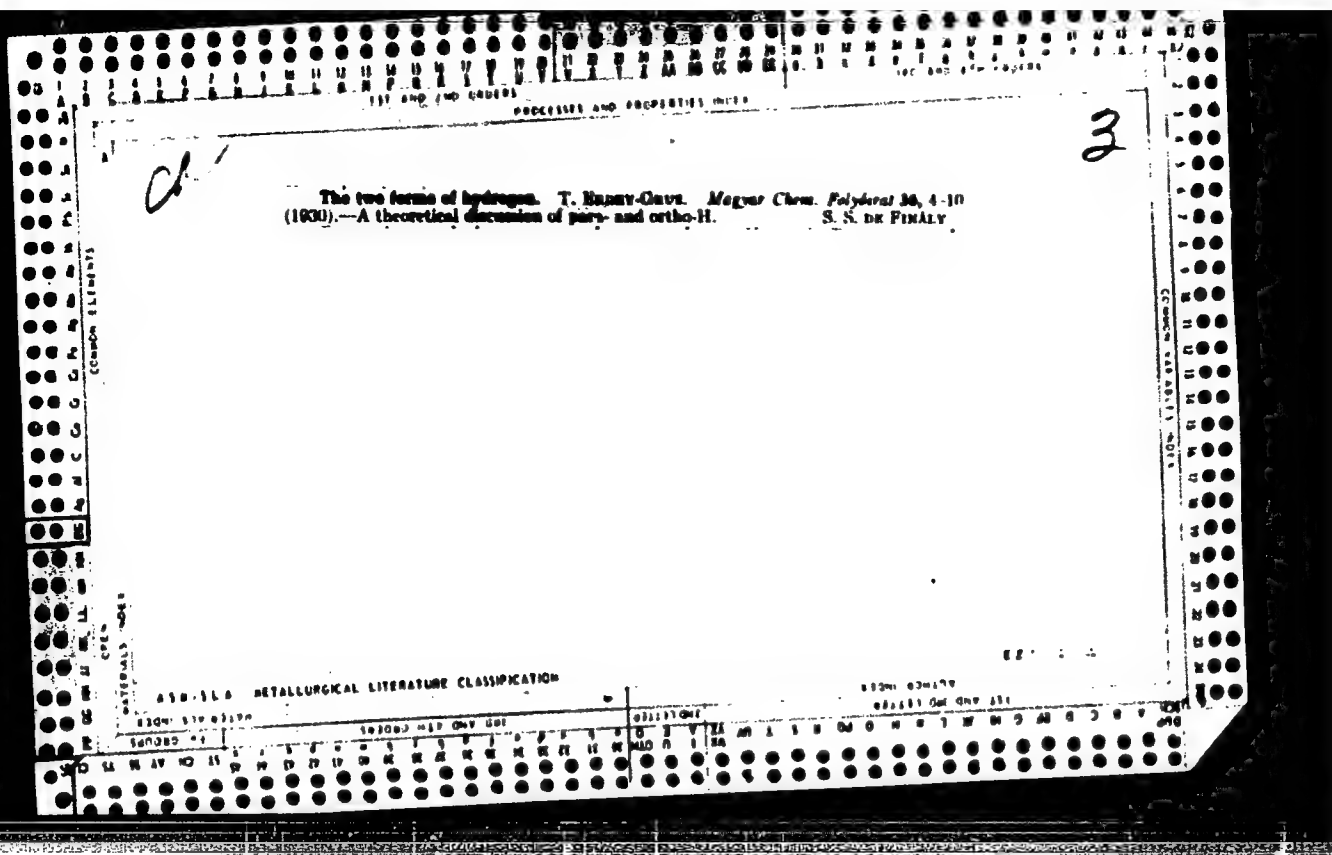
1. Problems of and experiences with coordinating the main task of the long-range research entitled "Investigation of the mechanism of chemical processes as well as the regularities of chemical industrial operations." Kem tud kozl MTA 20 no.2: 199-229 '63.

1. Magyar Tudományos Akadémia levelező tagja; "A Magyar Tudományos Akadémia Kémiai Tudományok Osztályának Közleményei" szerkesztő bizottsági tagja (for Szabo). 2. Veszpremi Vegyipari Egyetem rektora; "A Magyar Tudományos Akadémia Kémiai Tudományok Osztályának Közleményei" szerkesztő bizottsági tagja (for Polinszky). 3. Magyar Tudományos Akadémia Központi Kémiai Kutató Intézete igazgatóhelyettese (for Nagy). 4. Eötvös Loránd Tudományegyetem Fizikai Kémiai és Radiológiai Tanszéke. 5. Magyar Tudományos Akadémia Műszaki Kémiai Kutató Intézetének igazgatója; "A Magyar Tudományos Akadémia Kémiai Tudományok Osztályának Közleményei" szerkesztő bizottsági tagja (for Korach). 6. Akadémia Elektrokémiai Kutató Csoport vezetője; "A Magyar Tudományos Akadémia Kémiai Tudományok Osztályának Közleményei" szerkesztő bizottsági tagja (for Lengyel).
(cont. on next card)

ERDEY-GRUZ, Tibor, akadémikus

The state of natural sciences and the Hungarian national
long-range scientific research plan. Magyar Tud 70 no.1:7-18
Ja '63.

1. Tudományos és Felsőoktatási Tanács elnöke.



Control of chemical processes. Tilhor Erley-Gruz.
Termination. Academy 76, 421-33 (1937). Activity of
enzymes, catalysts and biocatalysts in chem. reactions
and biol. procedures is described. S. S. de Vinálly

117 AND 118 CROSS										119 AND 120 CROSS									
PROCESSING AND PROPERTY INDEX																			
<p>CA</p> <p>Differences of concentrations caused by diffusion and the effect of glass diaphragms on diffusion rate. Tibor Keleny-Gada, Antal Hunyik, Eva Popány, and Alajos Váli. <i>Hung. Acta Chim.</i> 1, No. 3, 7-26(1948).— Diffusion expts. were carried out with strong electrolytes (KCl, LiCl, HCl, NaCl, KBr, LiC₂H₃O₂, BaCl₂, and MgCl₂) in the presence of FrOH, sucrose, AcOH, propionic acid, butyric acid, crotonic acid, succinic acid, malic acid, citric acid and As₂O₃. A Jena glass diaphragm of type G 4 was used. The electrolyte was present at the beginning of the expt. only in the soln. on one side of the diaphragm. The concn. of the various added org. substances (excepting As₂O₃) seemed to grow in that part towards which the ions are diffusing. The quantity of transported substances is higher than the quantity of water transported by the diffusing ions at the same time with their hydrate sphere. This transport of org. substances is not a consequence of solvation of strong ions but seems to be connected with the mechanism of diffusion. The diffusion of electrolytes in presence of added org. substance is unavailable for the detn. of hydration values and makes doubtful the reliability of relative hydration values detd. by means of transport nos. The radius of the largest pores of diaphragms of type G 4 is 4.5-6.0 microns, 40-60% of the pores being not larger than 1.2-1.8 microns. Diaphragms of the type G 3 contain pores with a max. radius of 9-10 microns, about half of the pores being of the size 1.8-2.8 microns. The influence of diaphragms upon the rate of diffusion of KCl is approx. proportional to the total cross section of pores. The permeability of glass diaphragms for water and N₂ was also detd. The permeability and the effect of diminishing the rate of diffusion with most diaphragms seemed to be in a relation corresponding to their pore distribution.</p> <p style="text-align: right;">István Földi</p>																			
A18-51A METALLURGICAL LITERATURE CLASSIFICATION										CST-2-2-2-2-2									
FROM SYNDICATE										FROM BUREAU									
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CA

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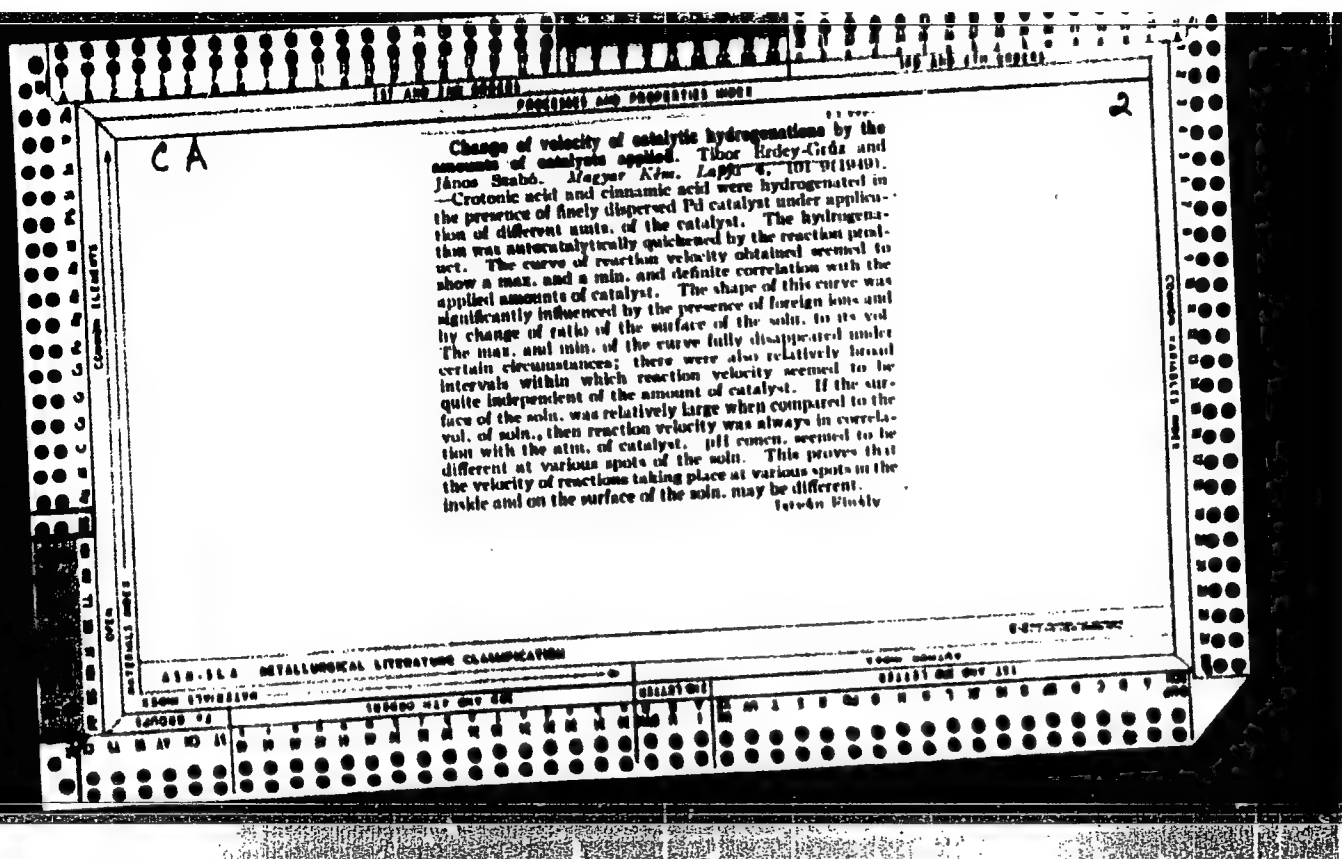
A special case of diffusion of two components in the same solution. Tihon-Medov-Nede-and Arslan Hunyar. *Hung. Acta Chim.* 1, No. 8, 27-32(1948). One part of a special diffusion app. was filled with a 1.0 N soln. of KCl contg. 8% propenol. The other part of the app. contained 8% propenol only, and both parts were sep'd. from each other by a glass diaphragm. Then the amounts of KCl and propenol passed through the diaphragm within 1, 2, and 4 days were detd. The quotient of these amts. seemed to diminish as the time passed. The quantity of 8% propenol soln. apparently transported by 1 mol. of the KCl soln. is the largest at the beginning of diffusion; it then gradually decreases owing to equilibration. The value for factor f obtained is 0.036. The measured value m_1/m_2 was 0.014 (1 day), 0.061 (2 days), and 0.024 (diffusion during 4 days).

István Flukly

ADDITIONAL LITERATURE CLASSIFICATION

100000 WITH SUB DIV

CLASSIFIED



C.A.

Electrolysis of complex silver salt solutions. Tibor Rády-Gárdi and Valéria Horváthy (Univ., Budapest, Hung.). *Magyar Kém. Lapja* 6, 834-31(1949).—A device was constructed for the purpose of scratching the surface of a silver electrode during electrolysis. This electrode was prep'd. from a thick wire by hammering it into a disklike shape. Solns. contg. various amts. of AgNO_3 , $\text{KAg}(\text{CN})_2$, and $\text{Ag}(\text{NH}_3)_2\text{OH}$ were electrolyzed and the mechanism of deposition of Ag on the cathode was studied. Gaseous N_2 was bubbled through the solns. during electrolysis to inhibit the dissolving of Ag in the cyanide soln. The max. current ds. were det'd. at which the Ag pptn. on the cathode still agreed with the law of Faraday; these values are called "limits" of 100% Ag pptn. (lim%). This limit was the highest in AgNO_3 solns. as compared to other solns. of identical concn., and the lowest in solns. of $\text{Ag}(\text{NH}_3)_2\text{OH}$. The value of lim% increased parallel to the increase of Ag concn., but showed no alteration when KNO_3 or NH_4OH was added to the soln. at a given Ag concn. The addn. of excess KCN to a soln. of $\text{KAg}(\text{CN})_2$ diminished the value of lim%. Increasing the temp. in the interval 0-20° increased the value of lim% by about 1% for each degree centigrade. A correlation of the anodic-dissolving effect to the current d., similar to that of the cathode was observed. The numeric value of lim% appeared somewhat higher for the anode. The presence of KNO_3 , KCN, and NH_4OH increased the value of anodic lim%. The results of expts. show no significant differences between electrolytic pptn. of Ag from Ag ion hydrates present in simple solns. of Ag salts and

electrolytic pptn. of Ag from complex ion solns. The mechanism of the process is the following: The ions are sepd. from the soln. by the force of the cathodic field, then they go to the surface of cathode, where they are neutralized. The Ag lost from the soln. by this pptn. is replaced by diffusion. The ion transfer due to the elec. current does not play a significant role in this respect. Calculs. based on these principles showed that the thickness of the layer between the surface of the cathode and the interior of the soln. must be about 10^{-3} cm. This is in accordance with results obtained in other fields. Diffusion actually transfers Ag in amts. corresponding to the cathodic current ds. up to the lim% values. In the case of current ds. above this rate, the diffusion is unable to replace the full amt. of Ag required. Existence of anodic dissolving effects below 100% is probably due to a layer covering the surface of the anode with solid salts. It seems that this layer forms when the velocity of Ag soln. becomes greater than the velocity of diffusion between the produced salt and that portion of the soln. which is in immediate contact with the surface of anode, and the latter thus becomes overatd. in respect to Ag compds.
I. Pindy

CA

4

Oscillographic analysis. The possibilities of a surface investigating method. Tibor Erdőy-Grúz (Univ., Budapest). *Magyar Kém. Folyóirat* 86, 83-7(1960).—The polarization capacity of a metallic surface depends on the condition of this surface and on the changes occurring thereon. The correlation of the polarization capacity with the changes in potential serves as a basis for detg. the conditions of the metallic surface and its changes, even when the surface area is unknown. For the detn. of polarization capacity the oscillographic method seemed to be suitable (*C.A.* 36, 1108). The oscillographic analysis of a Pt electrode in 1.0 N H₂SO₄, satd. with gaseous H₂ showed that the potential increase from 0 potential to H potential is not linear. A sharp increase at the beginning is later followed by a steadier period, and ends with another sharp increase. The const. value was 0.85–0.90 v. more pos. than the H-potential, corresponding to the potential of PtO, the test affirming the formation of an oxide layer on the surface of the Pt electrode. Other examples show the suitability of the oscillographic method for the study of surface processes of metals. The classic electrochem. law states that in the event of several possible electrode processes the process with the min. pos. or neg. potential will occur. Oscillographic studies proved that this law should be applied with caution, since under certain conditions, processes requiring more pos. potentials may occur. István Fényes

1951


000.715:020.191.2

98. The problem of aluminium corrosion in the chemical industry, by T. Erdey Grúz. („Magyar Kémikusok Lapja” -- Journal of the Hungarian Chemical Society -- Vol. V, No. 2, pp. 10-51, Feb., 1960.)

Metallic aluminium is much less resistant to corrosion than stainless steel, generally the circumstances may be chosen where corrosion of aluminium involves no danger from a technical or household point of view. Universal literature referring to the passivity and corrosion of aluminium alloys is dealt with; the effects of composition and the condition of various aluminium alloys on corrosion are discussed in detail. When applying aluminium in the chemical industry it must be kept in mind that aluminium

alloys with copper content generally have better mechanical properties but less resistance to corrosion, and that copperless aluminium alloys with iron content generally have poor mechanical properties. As stressed by recent experiments the significance of mechanical factors in the formation of the so-called intergranular corrosion will greatly influence the methods of mechanical treatments of aluminium alloys in the future. The influence of soldering and welding on the corrosion of aluminium is of extreme practical importance. It is very important that a perfectly smooth surface should precede all welding, since cavities or capillaries may act as sources of corrosion. The conditions of environment and the degree to which mechanical stresses are involved are also significant factors in the formation and development of corrosion. Practical methods for preventive treatments of aluminium corrosion are discussed, including structural designs which lessen the possibilities of corrosion, as well as inhibitors employable in the anti-corrosion treatment of aluminium and its alloys and various anti-corrosive films, dyes, and outdrying procedures. The chemical resistance of aluminium under given conditions is sufficiently high for extensive utilisation in the chemical industry, however, it must not be applied without a thorough investigation of the local operating conditions.

①
Oscillographic Analysis; Feasibility of a Method of Examining (Metal)
Surfaces (Oszcillografikus Analizis; Egy Felületvizsgálati Módszer
Lehetősége). Tibor Erdey-Gruz. (Magyar Kémiai Folyóirat, No. 2, 1950,
p. 83.) Gt. Brit., RAE Lib. Trans. 463, May, 1954. 10 pp. 16 refs.



181121

HUNGARY/Chemistry - Catalysts

"Dependence of the Rate of Hydrogenation on the Quantity of Catalyst." in German, J. Erdey-Gruz, J. Szabo, Inst Phys Chem and Radiol Budapest U

pp 46-55

Acta Chimica Acad Sci Hungaricae" Vol I, No 1,

Examined hydrogenation of crotonic and cinnamic acid in presence of finely divided palladium. Hydro- generation product has autocatalytic effect on reaction. Rate of hydrogenation as function of catalyst quantity has max and min. It is af- fected by concn of soln, presence of foreign

181122

HUNGARY/Chemistry - Catalysts (Contd)

ions, and changes in surface/vol ratio of so- lutions, and can even become independent of quantity of catalyst. If surface/vol ratio is large, hydrogenation will be proportional to quantity of catalyst.

ERDEY-GRUZ, TIBOR.

Elmeleti fizikai kemia (irtak) Erdey-Gruz Tibor es Schay Geza. 2. kiad. Budapest, Tankonyvkiado. (Egyetemi tankonyv) (Theoretical and physical chemistry; a university textbook. 2d ed. illus., diags., graphs, indexes, tables)
Vol. 1. 1955. 619 p.

SO: Monthly Index of East European Accession (EEAI) LC. Vol. 7, No. 5, 1958

ERDEY-GRUZ, T. (Budapest)

Hungary

T. Erdey-Gruz, author of "Influence of cations upon oxygen overvoltage," presented at the 4th ~~Electrochemical~~ Conference Conference, Moscow, 1-6 Octbber, 1956,
Electrochemical

SOURCE: Program to the 4th International Conference on Electrochemistry, Moscow, 1-6 October 1956, Unclassified.

ERDEY-GRUZ, T.

Category : USSR/General Problems - Problems of Teaching

A-3

Abs Jour : Ref Zhur - Fizika, No 2, 1957 No 2783

Author : Erdei-Gruz, Tibor

Title : Physics Teaching in Hungarian Schools

Orig Pub : Fizika v shkole, 1956, No 4, 51-53

Abstract : No abstract

Card : 1/1

ERDEV-GRUZ, T

19. The rate of catalytic hydrogenations

Grúz, K. Z. *Magyar Kémiai Folyóirat*
Vol. 62, 1956, No. 9, pp. 302--306, 21 figs.

The rate of hydrogenation of cinnamic acid sodium salt was studied in aqueous solution in the presence of palladium catalyst on barium sulphate carrier. The rate of hydrogenation as a function of the amount of the catalyst is influenced by several factors. Increase of the degree of dispersion of the catalyst increases the difference between the maximum and minimum reaction rate, and increases the rate of the reaction near the maximum. By increasing the surface-volume ratio of the solution, a parallel increase of the rate of hydrogenation is obtained. Presence of other electrolytes produces various effects: some of them (KCl, BaCl₂) decrease, others (K₂SO₄, HCl, NaOH) increase or decrease the reaction rate depending on their concentration. If ammonium salts are present the rate of hydrogenation is independent of the amount of catalyst within a wide range. Reducing the pressure of hydrogen causes an almost proportional decrease of the reaction rate. An interpretation of the empirical connection between hydrogenation rate and amount of catalyst is suggested.

ERDEY-GRUZ T.

B-13

HUNGARY/Physical Chemistry - Surface Phenomena. Adsorption.
Chromatography. Ion Exchange.

Abs Jour : Ref Zhur - Khimiya, No 8, 1958, 24356

Author : Erdey-Gruz T., Nagy F.

Inst : Hungarian Academy of Sciences.

Title : Adsorption of Ethylene at Activated Carbon in Water
Suspension.

Orig Pub : Acta chim. Acad. sci. hung., 1957, 12, No 1, 101-114

Abstract : Detailed description of a unit, designed by the authors,
for recording of adsorption isotherms (AI) by the volume-
tric method at a constant pressure, which is also suitable
for determination of AI and adsorption kinetics of
adsorbents suspended in water. Over the range of 0-500
mm Hg were recorded AI of ethylene at dry, activated wood
charcoal (I; 18.6, 20.0 and 25.0°) with a specific

1/2

Erdely, L.

Distr: H_2O

7
Oxidation-reduction titrations in nonaqueous media.
 Laszlo Erdely and Gyorgy Rády (Tech. Univ., Budapest).
~~Acta Chim. Acad. Sci. Hung.~~ 13, 81-93 (1958) (in German).
 —By using ascorbic acid solns. as the reductant it is possible
 to det. $Br_2/Au(III)$ and $Hg(II)$ by potentiometric titration
 in glacial $AcOH$. ICl , $KMnO_4/Cr(VI)$, and $V(V)$ can also
 be titrated by using ascorbic acid solns. as reductant but the
 stoichiometry of these reactions was not detd. The end
 points are indicated by reproducible potential changes in all
 cases, an abrupt change of 180 to 600 mv. being observed in
 those systems suitable for detn. Pt and satd. aq. calomel
 electrodes were used, and the diffusion potential was neg-
 lected.
 Mark M. Jones

10

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JH JD

L. ERdey

Distr: 4E2c/4E3d/4E2c(j

Decomposition of hydrogen peroxide in an alkaline solution in the presence of a copper citrate complex. L. ERdey and I. Inczédy (Tech. Univ., Budapest). *Acta Chim. Acad. Sci. Hung.* 17, 93-111 (1968) (in German). — A study of H_2O_2 decompn. in alk. soln. showed that on increasing pH values in the presence of a Cu^{++} -citrate complex the rate of decompn. rises linearly, whereas in pure alk. soln. there is a max. at pH 12. In the decompn. the homogeneous process is clearly discernible from the much slower process of O development on the walls. The HO_2 radical and intermediates of a brown Cu peroxy compd. (I) play a role in the homogeneous process. The rate of the over-all process is detd. by stationary I concns. and the concn. of perhydroxyl ions. When the initial mole ratio of H_2O_2 and Cu^{++} ions exceeds 100, the stationary concn. of I is stable at a larger interval, within which a 1st-order reaction is observed. The const. stationary concns. are approx. independent of pH and of H_2O_2 concn. The activation energy of decompn. is 12 kcal./mole. Perhydroxyl ions are dominant in the decompn. and their activity is reduced by undissocd. H_2O_2 mols. The homogeneous process is nearly independent of glass surface area in contact with the soln., but the rate of O development is proportional to surface area and, at high surface/vol. ratios, can approach the rate of decompn.

M. J. D. Low

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ERDEY-GRUBZ, I

ATCA CHIMICA
Academiae Scientiarum Hungaricae
Vol 13, Nrs 1-3, 1967

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THE ACTION OF CATIONS ON THE POLARISATION POTENTIAL OF PLATINUM
ANODES AND ON THE RATE OF HYDROGEN EVOLUTION

I. ERDEY-GRUBZ and I. SZABO
(Institute of Physical Chemistry, Hungarian Academy of Sciences, Budapest)

Summary

37
1. When a platinum electrode immersed in a solution of a substance which is anodically polarised at a low current density, the anodic polarisation potential decreases with the logarithm of current density (i).

$\log i = 4 - 0.11 \log i$

the electrolyte concentration. The anodic polarisation potential can be detected even after a lengthy electrochemical reduction of the electrode. After evolution from water, the rate of evolution is not affected.

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1. On raising the current density above 10^{-2} – 10^{-1} amp./sq. cm (at an anode potential of about $\varphi = +2.0$ V), the polarisation potential rapidly increases, and the course of reaction alters in that persulphates are gradually formed begins in solutions of higher concentration.
2. In pure solutions of sulphuric acid of 2.0 N and above this concentration, the polarisation potential of the anode increases at a low current density approximately linear with the concentration. Parallel to the increase of polarisation potential, the value of b rises from 0.106 (in a 2.0 N solution) to 0.139 (in a 9.0 N solution of sulphuric acid).
3. When the sulphates of different metals are dissolved in a solution of sulphuric acid, the polarisation potential of the anode becomes more positive (compared at an identical current density) even at an unchanged total concentration of electrolyte or at an unchanged concentration of sulphuric acid. Referring to their action on increasing polarisation, the sequence of the metal cations examined proved to be as follows: $K^+ > Al^{3+} > NH_4^+ > Zn^{2+} > Na^+ > Mg^{2+} > Li^+$. By rising polarisation potentials, also the value of b increases from 0.106 to 0.135 in a 2.0 N solution. Metal cations showed similar action in a 9.0 N solution of sulphuric acid.
4. In solutions of pure salts of the metals examined, the correlation of the change of polarisation potential with the nature of the cation was similar to those observed in the presence of sulphuric acid.
5. K^+ ions, with the increase of polarisation potentials, showed an action of identical character both in the case of smooth and platinated platinum electrodes, and on nickel electrodes.

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6. The action of increasing polarisation (measured at a low constant current intensity) showed an approximately linear correlation with the logarithm of the concentration of K^+ .
7. To interpret the increase of polarisation potentials caused by metal cations, the authors presume that metal cations are adsorbed in the diffuse portion of the double layer, by the ions SO_4^{2-} and HSO_4^- , which are directly adsorbed by the electrode surface. Metal cations, bound this way, alter the linkage of water molecules adsorbed by the electrode surface with other water molecules, and deform the latter in that that the activation energy of the electron leap from the water molecules to the electrode rises. Consequently, the electrode process becomes slower and, respectively, higher polarisation potential is required to maintain a given rate (current density).

ERDEY GRUZ, I.
 ACTA CHIMICA
 Academiae Scientiarum Hungaricae
 Vol 13, Nrs 1-2, 1957

EFFECT OF ALTERNATING CURRENT ON THE OVERPOTENTIAL OF OXYGEN
 ON PLATINUM ANODE IN SOLUTIONS OF SULPHURIC ACID

I. ERDEY GRUZ and J. SAFARIK

(Institute of Physical Chemistry, L. Eötvös University, Budapest)

Received December 2, 1956

Summary

The changes of the polarisation potential of a platinum anode (against a normal hydrogen electrode of 0 v.) under the action of alternating current superimposed to direct current, were examined in a 3.0 N and 9.0 N solution of sulphuric acid. When the frequency of the alternating current was observed, the strength of which was varied, the overpotential of oxygen evolution decreased. The current density of direct current polarisation is approximately 10 mA/cm² in sulphuric acid.

The polarisation curves ($\eta = A + b \log i$) proved to be linear with few exceptions, when working in the interval of oxygen evolution and at low current densities (e.g. 1-4). In a 3.0 N solution the value of b slightly increases under the action of alternating current. In a 9.0 N solution, however, under the action of alternating current of 50-1000 Hz, the value of b significantly decreases. The value of b did not appreciably change (about 10%) when the frequency of alternating current was varied with direct current. When the frequency of alternating current was varied with direct current, the value of b did not appreciably change (about 10%).

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Alternating current of low frequency reduced polarisation in a 2.0 N solution of sulphuric acid by about 0.05 v. (Fig. 5, Table I), the minimum value of polarisation (and the maximum of depolarisation) appears at about 500-1000 Hz. Further rise in frequency is followed by a slight increase in polarisation. The use of longer electrodes (10 cm) leads to a more pronounced effect.

From these data it follows that the alternating current of low frequency (500-1000 Hz) reduces the polarisation of a 2.0 N solution of sulphuric acid by about 0.05 v.

On the basis of the observed phenomena, it can be stated that the overpotential of oxygen evolution is reduced by a superposed alternating current. In other words, the alternating current promotes the process of oxygen evolution.

On the basis of the observed phenomena, it can be stated that the overpotential of oxygen evolution is reduced by a superposed alternating current. In other words, the alternating current promotes the process of oxygen evolution. The formation of peroxide is also promoted by the alternating current. The rate of oxygen evolution is faster than that of hydrogen evolution.

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below that of reductions. However, at higher frequencies, the degree of reduction of alternating current decreases, due to the limited value of the rate of reduction of higher frequencies. The rate of reduction of higher frequencies is not hydrolysis, respectively, the rate of reduction of higher frequencies is not hydrolysis.

The experiments show that the rate of reduction is the rate of formation of products.

~~SECRET~~ ERDEY-GRUZE, Tibor

Distr: 4E43 7
/ Speed of catalytic hydrogenation. Tibor Erdely-Grúz
and Károly Zimmer (Bolyai Loránd Univ., Budapest,
Hung.). Magyar Kém. Folyóirat 62, 302-8 (1967).—Rates
of hydrogenation of Na cinchonate were measured in aq.
soln. in the presence of Pd catalyst on BaSO₄ carrier. A
great number of assumptions had to be made when attempt-
ing to explain the empirical correlation between rate and
amt. of catalyst. Francis J. Schmidt

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J. J. J.

ERDEY-GRUZ, TIBOR

The effect of cations on the polarization potential of platinum anodes on the oxygen overvoltage. Tibor Erdey-Grúz and Imre Szafarik (Eötvös Loránd Univ., Budapest). Magyar Kém. Folyóirat 63, 221-4 (1967). The mechanism of the processes at the Pt anode in H_2SO_4 solns. of different concns. was investigated by studying the effects of Li^+ , Na^+ , K^+ , NH_4^+ , Mg^{2+} , Zn^{2+} , Al^{3+} ions on the polarization potential. The presence of metallic ions in the H_2SO_4 solns. resulted in increased polarization at the Pt anode and in increased slope of the Tafel lines. The same effect of K^+ ions was observed at platinized Pt and Ni electrodes. The effect on the anodic polarization was directly proportional (at const. c.d.) to the logarithm of the K^+ concn. The behavior of the cations was explained by adsorption in the diffuse part of the double layer increasing the activation energy of the electrolytic O evolution. Saul Patai

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Jan

ERDEY-GRÜZ, TIBOR

B-12

HUNGARY/Physical Chemistry - Electrochemistry.

Abs Jour : Ref Zhur - Khimiya, No 7, 1958, 20777

Author : Tibor Erdey-Grüz, Imre Safarik.

Inst : -

Title : Influence of Alternating Current on Oxygen Overvoltage on Platinum Electrode in Sulfuric Acid Solution.

Orig Pub : Magyar kem. folyóirat, 1957, 63, No 9, 237-242

Abstract : The dependence of oxygen overvoltage on smooth Pt at i from 10^{-5} to 1 amp. per sq.cm in 2 n. H_2SO_4 , as well as 9 n. H_2SO_4 on the frequency (ν) of an additionally superimposed alternating current (50 to 20,000 cycles) was studied. With the rise of ν , the potential of Pt passes (at a given i) through a minimum, the depth of which at $i = 10^{-5}$ to 10^{-2} amp. per sq.cm is 0.05 to 0.06 v in the case of 2 n. H_2SO_4 , and 0.32 to 0.36 v in the case of 9 n. H_2SO_4 , and at $i = 1$ amp. per sq.cm it drops to 0.003 v and zero correspondingly. The depolarizing action of the

Card 1/2

HUNGARY/Physical Chemistry. Electrochemistry.

B

Abs Jour: Ref Zhur-Khin., No 15, 1958, 49711.

In solutions of KOH and KCl the minimum of (Λ, N) curves is flattened, is located in the interval $N = 40-60\%$, and corresponds to the maximum of the (Λ, N) curve. With increasing N up to $80-90\%$ the temperature coefficient (TC) of Λ of all three electrolytes increases, while at higher N -- it drops. TC of Λ , over a wide range of N , varies approximately linearly with increasing N , and only with very low contents of water and I the TC decreases sharply. Energy of activation of electric conductivity A (in kcal/mole in all instances) is about 2.3 in pure I . With increasing N up to $75-85\%$ A increases, after that it drops. Maximum values of A : 4.6 for HCl, 5.1 for KOH, 4.6 for KCl. For Λ the A values increase with increasing N ,

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HUNGARY / Physical Chemistry. Electrochemistry.

B

Abs Jour : Ref Zhur - Khimiya, No 12, 1959, No. 41723

Author : Erdey-Gruz, T.; Majthenyi, L.

Inst : Hungarian AS

Title : The Transfer Mechanism of Hydrogen and Hydroxyl Ions. II. Transfer Numbers of HCl, KOH, KF and KCl in Water-Methanol Mixtures at Temperatures Ranging from 5-25°.

Orig Pub : Acta chim. Acad. scient. hung., 1958, 16, No 4, 417-438

Abstract : The transfer numbers n for HCl, KOH, KF and KCl were determined by the moving boundary method from a mixture of methanol (I) and water at temperatures ranging

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HUNGARY / Physical Chemistry. Electrochemistry.

B

Abs Jour : Ref Zhur - Khimiya, No 12, 1959, No. 41723

raised. From the electroconductivity data published earlier (R. Zh. Khim, 1959, No 5, 14759), and the values of n obtained, the ion mobility μ , expression $\mu \cdot \eta$ (η - viscosity), and temperature coefficients of the mobility (TCM) were calculated. TCM dependence on the solvent's water content passes through a maximum for all ions. The greatest maximum was observed with F^- and the smallest for H^+ ions. Maxima on TCM-composition, and η -composition curves were observed at identical compositions for H_3O^+ and OH^- ions, while those for K^+ , Cl^- and F^- occurred at different compositions. K^+ ion mobility was the same in KOH, KF and KCl solutions,

Card 3/4

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HUNGARY/Physical Chemistry. Electrochemistry.

Abs Jour: Ref Zhur-Khin., No 5, 1959, 14759.

Author : Erdey-Gruz T., Hajthenyi L.

Inst : ~~Central Research Institute for Chemistry~~

Title : The Mechanism of Movement of Hydrogen and Hydroxyl Ions. II. The Transfer Numbers of HCl, KOH, KCl and KF in Methanol-Water Mixtures at 5 and 25°.

Orig Pub: Magyar kem. folyoirat, 1958, 64, No 6, 212-220.

Abstract: The transfer numbers (TN) of HCl, KOH, KCl and KF in methanol (I) - water mixtures of various compositions have been measured at 5 and 25°. The TN in the case of HCl and KOH strongly depend on the water content in the mixture; in the case of KCl and KF - they are insignificant. The TN of H⁺,

Card : 1/3

27

ERDEY-GRUZ, T., MAJHENYI, L.

Mechanism of migration of hydrogen and hydroxyl ions. V. Effects of the composition of ethanol and water on the transference numbers and ion mobilities of dissolved LCL, KOH, KF AND KCl at 5 and 25 C. In German, p. 73

ACTA CHIMICA. Budapest, Hungary, Vol. 20, No. 1, 1959

Monthly List of East European Accessions (EEAI) LC, Vol. 9, No. 2 Feb. 1960
Uncl.

ERDEY-GRUZ, T.; MAJTHENYI, L.

Mechanism of migration of hydrogen and hydroxyl ions. VI. Effect of the temperature and composition of glycol-water mixture on the transference numbers and ion mobilities of dissolved HCl, KOH, KF and FCl at 5° and 25° C.

ACTA CHIMICA. (Magyar Tudományos Akadémia) Budapest, Hungary. Vol. 20
No. 2, 1959

Monthly Lists of East European Accessions, (EEAI) LC, Vol. 9, No. 1, 1960

Uncl.

ERDEY-GRUZ, Tibor; KUGLER, Elvira; HIDVEGI, Judit

Migration mechanism of hydrogen and hydroxyl ions. Pt. 3. Magyar
kém folyoir 65 no.3:114-123 Mr '59.

1. Eotvos Lorand Tudományegyetem Fizikai-Kémiai Tanszéke, Budapest.
2. "Magyar Kémiai Folyoirat" felelős szerkesztője (for Erdey-Gruz).
3. "Magyar Kémiai Folyoirat" szerkesztősegi titkára (for Kugler).

ERDEY-GRUZ, T.; KUGLER, E.; HIDVEGI, J.

Mechanism of the migration of the hydrogen and hydroxyl ions. IV. Effect of the constitution of glycol-water mixtures on the conductivity of dissolved hydrochloric acid, potassium hydroxide, potassium fluoride, and potassium chloride as well as their viscosity at 5° and 25°C. p. 152.

MAGYAR KEMIAI FOLYOIRAT. Budapest, Hungary. Vol. 65, no. 4, Apr. 1959

Monthly List of East European Accessions (EEAI), LC. Vol. 8, No. 9, September 1959
Uncl.

ERDEY-GRUZ, Tibor, r.tag (Budapest)

An account made by the Section's leadership; also, remarks by Gyula Hardy and others. Kem tud kozl MTA 14 no.2:141-175 '60. (EEAI 10:2)

1. Osztalytitkar, Magyar Tudomanyos Akademia Kemiai Tudomanyok
Osztalya, Budapest.
(Hungarian Academy of Sciences) (Hungary--Chemistry)

ERDEY-GRUZ, Tibor, akademikus (Budapest)

Transition of quantitative changes into qualitative ones as seen by a
chemist. Magyar Tud. 67 no.8:467-484 Ag '60. (EEAI 9:11)
(Chemistry)

ERDEY-GRUZ, Tibor, akademikus (Budapest)

Quantitative changes turning into qualitative ones, as seen by the
chemist. II. Magyar tud 67 no.9:529-543 S '60. (EEAI 9:12)
(Chemistry) (Gases) (Heat)
(Molecules) (Solutions)

ERDEY-GRUZ, Tibor, akademikus (Budapest)

High-level and up-to-date teaching in the institutions of higher
education. Magy tud 67 no.12:711-715 D '60. (EEAI 10:3)
(Hungary--Universities and colleges)

ERDEY-GRUZ, Tibor, egyetemi tanár (Budapest)

Modern science and religion; the voice of natural sciences. Munka 11
no.1:20-21 Ja '61.

(Religion and science)

ERDEY-GRUZ, Tibor, akadémikus (Budapest); CHOLNOKY, László; SZABÓ, Zoltán;
SZÉKER, Gyula, kandidátus; FOLDI, Zoltán; LANGYEL, Sándor, a tudományok
doktora; TAKÁCS, Pál, kandidátus

An account of the 1960 work of the Section of Chemical Sciences,
Hungarian Academy of Sciences. Kem tud közl MTA 15 no.4:401-460 '61.

1. Osztálytitkár, Magyar Tudományos Akadémia Kémiai Tudományok Osztálya,
Budapest és Szerkesztő, Magyar Tudományos Akadémia Kémiai Tudományok
Osztályának Közleményei (for Erdy-Gruz) 2. Lev. tag, Magyar Tudományos
Akadémia Kémiai Tudományok Osztályának Közleményei (for Chólnoky, Szabó,
Foldi) 3. Szerkesztőbizottsági tag, Magyar Tudományos Akadémia Kémiai
Tudományok Osztályának Közleményei (for Lengyel)

(Hungarian Academy of Sciences) (Hungary—Chemistry)

ERDEY-GRUZ, Tibor, dr.

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